A population-level prediction tool for the incidence of first-episode psychosis: translational epidemiology based on cross-sectional data


This version is available from Sussex Research Online: http://sro.sussex.ac.uk/id/eprint/62397/

This document is made available in accordance with publisher policies and may differ from the published version or from the version of record. If you wish to cite this item you are advised to consult the publisher’s version. Please see the URL above for details on accessing the published version.

Copyright and reuse:
Sussex Research Online is a digital repository of the research output of the University.

Copyright and all moral rights to the version of the paper presented here belong to the individual author(s) and/or other copyright owners. To the extent reasonable and practicable, the material made available in SRO has been checked for eligibility before being made available.

Copies of full text items generally can be reproduced, displayed or performed and given to third parties in any format or medium for personal research or study, educational, or not-for-profit purposes without prior permission or charge, provided that the authors, title and full bibliographic details are credited, a hyperlink and/or URL is given for the original metadata page and the content is not changed in any way.
A population-level prediction tool for the incidence of first-episode psychosis: translational epidemiology based on cross-sectional data

James B Kirkbride,1 Daniel Jackson,2 Jesus Perez,3 David Fowler,4 Francis Winton,5 Jeremy W Coid,6 Robin M Murray,7 Peter B Jones1,8

ABSTRACT

Objectives: Specialist early intervention services (EIS) for people aged 14–35 years with first episodes of psychosis (FEP) have been commissioned throughout England since 2001. A single estimate of population need was used everywhere, but true incidence varies enormously according to sociodemographic factors. We sought to develop a realistically complex, population-based prediction tool for FEP, based on precise estimates of epidemiological risk.

Design and participants: Data from 1037 participants in two cross-sectional population-based FEP studies were fitted to several negative binomial regression models to estimate risk coefficients across combinations of different sociodemographic and socioenvironmental factors. We applied these coefficients to the population at-risk of a third, socioeconomically different region to predict expected caseload over 2.5 years, where the observed rates of ICD-10 F10-39 FEP had been concurrently ascertained via EIS.

Setting: Empirical population-based epidemiological data from London, Nottingham and Bristol predicted counts in the population at-risk in the East Anglia region of England.

Main outcome measures: Observed counts were compared with predicted counts (with 95% prediction intervals (PI)) at EIS and local authority district (LAD) levels in East Anglia to establish the predictive validity of each model.

Results: A model with age, sex, ethnicity and population density performed most strongly, predicting 508 FEP participants in EIS in East Anglia (95% PI 459, 559), compared with 522 observed participants. This model predicted correctly in 5/6 EIS and 19/21 LADs. All models performed better than the current gold standard for EIS commissioning in England (716 cases; 95% PI 664–769).

Conclusions: We have developed a prediction tool for the incidence of psychotic disorders in England and Wales, made freely available online (http://www.psymaptic.org), to provide healthcare commissioners with accurate forecasts of FEP based on robust epidemiology and anticipated local population need. The initial assessment of some people who do not require subsequent EIS care means additional service resources, not addressed here, will be required.

BACKGROUND

Commissioners of health and social care require precise information on the health needs of their local populations,1 especially if parity of mental and physical health is to be realised.2 Mental health disorders alone represent the leading disease burden in the UK.
Psychosis incidence prediction

ARTICLE SUMMARY

Strengths and limitations of this study
- Our modelling approach used robust epidemiological data from two large studies of first episode psychosis in England to provide estimates of incidence in a third study region, producing accurate FEP forecasts.
- While our models provide estimates of the expected clinical burden of FEP in the community, services may see a broader range of psychopathology consuming resources, or incepted rates may be influenced by supply-side organisational factors.
- Owing to data availability, it was not possible to validate our prediction tool in settings outside of England and Wales, or for specific psychotic disorders. As data become available, we will extend the capability of our prediction tool, including into other settings and disorders.

(22.8%). They contribute substantially to healthcare expenditure and societal costs even before physical ill health is taken into account. The Centre for Mental Health estimated the total costs of mental health to British health services and society at £105 billion in 2009/2010, a figure expected to double over the next 20 years. These are serious challenges compounded by a paucity of information on which to commission appropriate services. Early intervention in psychosis services (EIS) for people aged 14–35 years with a first episode of psychosis (FEP) offer a useful example of failure to map services to local need.

EIS are a major evidence-based innovation, systematically commissioned throughout England and Wales over the past decade. When EIS intervention is sustained, there is evidence that people with psychosis achieve better functional and social outcomes. Such services are also highly cost-effective. However, EIS were originally commissioned on an anticipated rate of 150 new cases of any psychotic disorder per 1 000 000 of the total population per year in the Department of Health’s Mental Health Policy Implementation Guide (MH-PIG). In 2001 in England and Wales, 29.3% of the population were aged 14–35 years, meaning that the MH-PIG commissioned incidence rate was approximately 51 cases/100 000 person-years in the age range covered by EIS. Following their deployment, anecdotal reports began to emerge from EIS in different regions to suggest that a uniform figure for commissioning was simultaneously underestimating and overestimating the actual observed need in urban and rural populations, respectively. Recent epidemiological evidence of FEP incidence in rural communities in England has suggested that rates are somewhat lower than the uniform figure upon which services were commissioned, confirming previous calls that a ‘one-size-fits-all’ prescription for EIS implementation is unlikely to lead to the efficient allocation of finite mental health resources.

Using rich epidemiological data on variation in the incidence of FEP according to major sociodemographic risk factors, we describe the development and validation of a population-level prediction tool capable of accurately estimating the expected incidence of psychiatric disorder, based on the sociodemographic structure of the population in a given region. Applied to FEP as proof-of-concept, we show that it is possible to closely predict the expected incidence in a given population, where the observed count of cases was within the prediction intervals (PI) forecast by our models. We applied our most precise prediction model to the population of England and Wales to provide health commissioners with a translational epidemiological prediction tool to underpin information-based service planning.

METHODS

Our prediction models were based on epidemiological data from the Aetiology and Ethnicity in Schizophrenia and Other Psychoses (ÆSOP) and the East London First Episode Psychoses (ELFEP) studies, two methodologically similar population-based FEP studies. We fitted various count-based regression models with different combinations of sociodemographic and socioenvironmental factors, well established in the literature to be associated with the incidence of psychotic disorder. We first established the relative apparent validity of each model by estimating model-fit diagnostics to assess how well each model fitted the empirical data (henceforth, the prediction sample). We next sought to estimate the external validity of each model by applying model-based parameter coefficients to the population structure of a purposefully different region of England, East Anglia (henceforth, the validation sample). This out-of-sample prediction technique allowed us to obtain the expected incidence of disorder in this region forecast by each model, which we compared with observed rates simultaneously ascertained in this region via the ongoing Social Epidemiology of Psychoses in East Anglia (SEPEA) study. We performed various model-fit diagnostics to identify which, if any, model demonstrated utilisable predictive capability.

Empirical data underlying prediction models (prediction sample)

Case ascertainment (numerator)
The designs of the ÆSOP and ELFEP studies have been described in detail elsewhere, with features relevant to the present paper summarised here. Case ascertainment took place over 2 years in ELFEP (Newham: 1996–1998; Tower Hamlets & Hackney: 1998–2000) and the Southeast London and Nottingham centres of the ÆSOP study (1997–1999), and over the first 9 months of 1997 in Bristol (ÆSOP). All service bases were screened regularly for potential new contacts aged 16–64 years (18–64 in ELFEP) resident within these catchment areas. Leakage studies were conducted to identify participants missed by this initial screen, but meeting inclusion criteria for FEP. All participants who received an ICD-10 F10–39 diagnosis for psychotic disorder
following assessment via the Schedules for Clinical Assessment in Neuropsychiatry were included in the incident sample, except those with an organic medical basis to their disorder or profound learning difficulty. Data on age-at-contact, sex and ethnicity were collected on included participants. We geocoded participants’ residential postcode at first contact to their corresponding local authority district (LAD) to allow us to model possible neighbourhood effects associated with the incidence of psychotic disorder, such as population density or socioeconomic deprivation.

Population at-risk
We estimated the population at-risk using the 2001 Census of Great Britain, adjusted for study duration, and stratified by age group (16–17, 18–19, then 5-year age bands), sex and ethnicity. Ethnicity was based on self-ascription according to 1 of 10 categories derived from the census: white British, non-British white, black Caribbean, black African, Indian, Pakistani, Bangladeshi, mixed white and black Caribbean, other mixed ethnic backgrounds and all other ethnicities.

Socioenvironmental variable estimation
We estimated LAD-level deprivation using the 2004 Index of Multiple Deprivation (IMD) in England, which estimated domains of deprivation using measures predominantly collected close to the time of our case ascertainment periods (see table 1).23 We z-standardised English LAD IMD scores to have a mean of zero and SD of 1, and extracted IMD z-scores for the 14 LADs in the ESOP and ELFEP studies. To inspect whether any particular deprivation domain was a better predictor of psychosis incidence than IMD, we also considered LAD-level income deprivation, employment deprivation and the extent of deprivation in our models (table 1). We estimated population density by dividing each LAD’s usual resident population by its area (in hectares), using ArcGIS V9.3 software.

Table 1 Description of included socioenvironmental variables†

<table>
<thead>
<tr>
<th>Variable</th>
<th>Classification and description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Multiple deprivation</td>
<td>Weighted data from routine national sources across seven domains: income, employment, education, health, barriers to housing and services, living environment, crime. Continuous, z-standardised scores for analysis</td>
</tr>
<tr>
<td>Extent of deprivation</td>
<td>Proportion of LAD population living in 20% most deprived SOA in England (%)</td>
</tr>
<tr>
<td>Income deprivation</td>
<td>Proportion of all people in LAD classified as income deprived (%)</td>
</tr>
<tr>
<td>Employment deprivation</td>
<td>Proportion of adults of working age in LAD classified as employment deprived (%)</td>
</tr>
<tr>
<td>Population density</td>
<td>Population density at LAD level (people per hectare)</td>
</tr>
</tbody>
</table>

†Validation sample sources: Population density—2001 mid-year census estimates; deprivation variables: 2010 Indices of Deprivation, predominantly collected from data sources just prior to the SEPEA case ascertainment period (2008); IMD, Index of Multiple Deprivation; LAD, local authority district; SOA, super output area.

Observed data for external validation of prediction models (validation sample)
Observed participants and population at-risk data for our validation sample were obtained from the SEPEA study, an ongoing study of the incidence of psychotic disorders incepted over 3.5 years (2009–2013) through one of six EIS covering 20 LADs and a subsection of 1 LAD (the town of Royston, Hertfordshire) in Norfolk (three EIS: West Norfolk, Central Norfolk, and Great Yarmouth and Waveney), Suffolk (one EIS) and Cambridgeshire, Royston and Peterborough (CAMEO North and South EIS).13

Case ascertainment
To establish the incepted incidence of FEP as seen through EIS, entry criteria for the SEPEA study were:

- Referral to an EIS in East Anglia for a suspected first episode of psychosis;
- Aged 16–35 years at first referral to EIS (17–35 years in CAMEO services);
- Resident within the catchment area at first referral;

At 6 months after EIS acceptance, or discharge from the service, whichever was sooner, we asked the clinician responsible for care to provide an ICD-10 F10–39 psychiatric diagnosis using all information available. We excluded participants without a clinical FEP diagnosis, or participants presenting with an organic basis to their disorder or profound learning disability. For the remaining participants, basic sociodemographic and postcode information was recorded and classified in the same way as in the prediction sample. We included participants presenting to EIS during the first 2.5 years of the ongoing SEPEA study.

Population at-risk
We estimated the population at-risk of East Anglia using 2009 mid-year census estimates published by the Office for National Statistics (ONS) at the LAD level, by age group, sex and ethnicity.24 These estimates used the 2001 census base, adjusted for immigration, births and deaths each year. It was not possible to obtain 2009 mid-year estimates for the town of Royston, because data were only published at LAD level. Here, we used denominator data from the 2001 census data in order to

estimate the population at-risk in Royston. We do not believe that this would have substantially invalid our results as this town represented 0.6% of the overall population at-risk (n=9555) in the SEPEA study. Denominator data were multiplied by 2.5 to account for person-years of exposure in the validation sample.

Socioenvironmental variable estimation
For each LAD in the SEPEA study, we obtained corresponding socioenvironmental variables to those included in our prediction sample, using updated data collected as close to the SEPEA case ascertainment period as possible. Population density was estimated using 2009 mid-term population estimates. Our measures of deprivation were derived from IMD 2010,25 which was estimated in an analogous way to 2004 data, but collected from sources obtained immediately prior to the SEPEA study.

Statistical techniques
Dataset generation
We constructed a dataset for the regression analysis of count data by pooling data from the AESOP and ELFEP studies (the prediction sample). Data were stratified by age group, sex, ethnicity and LAD, such that each stratum (N=2536) represented the total count of FEP cases in a unique sociodemographic group for a given LAD, with a corresponding estimate of the population at-risk, treated as an offset in our models. Our socioenvironmental measures (population density, deprivation) were adjoined to the dataset for each LAD. Population at-risk data from the validation sample were stratified in the same way and retained in a separate database. Here, the count of cases, which we wished to predict, was entered as a vector of missing data which would be populated with predicted case estimates following prediction modelling.

Prediction models
We used the prediction sample data to fit negative binomial regression models to obtain parameter coefficients of incidence for the sociodemographic and socioenvironmental factors included in each model. We considered the internal and external predictive capabilities of six models, all of which contained age group, sex, an age–sex interaction term and ethnicity. Model 1 contained no further covariates. Model 2 also included IMD. We replaced IMD with either income, employment or the extent of deprivation, respectively, in models 3–5. Model 6 included population density. Initial exploration of the prediction sample data indicated the presence of possible overdispersion (variance (δ^2=1.37) exceeded mean (μ=0.4) count of cases), so negative binomial regression was preferred to Poisson regression since it explicitly models any overdispersion with an extra dispersion parameter.

Apparent model validity and prediction
We assessed apparent model validity in three ways. First, we used Akaike’s Information Criterion (AIC) to assess the respective overall fit of each model to the data. Second, we conducted K-fold cross-validation to assess each model’s apparent validity to predict cases within the prediction sample. This method randomly allocated strata in the prediction sample into K subsets. Each model was then re-estimated on K–1 subsets (the training data) to predict the expected counts of cases in the Kth subset (the test data). This was repeated over K trials, such that each stratum in the dataset appeared exactly once as the test data. At the end of this process, we derived Lin’s concordance correlation coefficient (CCC) and 95% CI to estimate the correlation between the predicted and observed counts of cases across all strata in the prediction sample. Finally, we estimated the root mean squared error (RMSE) to determine the average error between fitted and observed values from each model. Lower RMSE scores indicated a smaller prediction error. The RMSE is derived as

$$\text{RMSE} = \sqrt{\frac{\sum_{i=1}^{n} (\gamma_i - \hat{\gamma}_i)^2}{n}}$$

where γ_i and ŷ_i are the observed and predicted counts of cases in the ith stratum, respectively, and n is the number of strata.

We repeated K-fold cross-validation h times, generating K new random divisions of the data each time. We retained model-fit diagnostics across Kh iterations, and reported the mean of Lin’s CCC and RMSE to provide summary cross-validation statistics for each model. We specified K=10 and h=20, as recommended for cross-validation to obtain precise model-fit diagnostics.26

External model prediction and validation
We retained parameter coefficients from each model (using the full prediction sample data) and applied these to the corresponding population at-risk in the validation sample dataset. This gave out-of-sample prediction estimates for the expected count of cases in each stratum of the validation sample, given the model. We summed expected counts across relevant strata to estimate the (1) total predicted count of cases in the SEPEA region, (2) predicted counts in each EIS and (3) predicted counts by LAD. These counts were further stratified by broad age group (16–35, 36–64 and 16–64 years). Because the census (denominator) data were unavailable for 35-year-olds alone (needed to estimate their contribution to predicted counts in the age range for EIS, 16–35 years), we assumed that the risk coefficient was the same across all ages within the 35–39-year-old age group. We apportioned predicted counts on a 1:4 ratio (35:36–39 years) to their respective broad age groups.

To determine how well the MH-PIG® figure of 51 new cases per 100 000 person-years for EIS performed as a
predictive tool, we also estimated the predicted count of cases in the validation sample under this scenario, which we termed ‘Model 7’.

We derived 95% PIs for all summary predictions from first principles, since their derivation is not straightforward, nor routinely implemented by statistical software. PIs are similar to CIs, but account for SEs introduced in both the prediction and validation samples. We developed a bootstrap-like approach to obtain PIs from each model by simulating 1000 model-based realisations of the quantities we wished to predict, where we took the parameters to be the maximum likelihood estimates. We obtained the lower and upper bounds of the PIs as the corresponding quantiles of the simulated realisations (see appendix for full details).

To assess each model’s external predictive capabilities, we considered five markers of predictive accuracy. We compared the number of times the observed count of cases in the SEPEA study fell within the PIs estimated from each model for (1) the SEPEA region, (2) at the EIS level and (3) at the LAD level. We also derived EIS-level (4) and LAD-level (5) RMSE scores to estimate prediction error from each model in our validation sample. We ranked model performance (1: best and 7: worst) on these five measures, and estimated an overall mean rank to determine the overall predictive validity of each model.

Observational data on first episode psychosis in our validation sample were not available for the age range 36–64 years, so external validation was restricted to the 16–35 year old age range. For completeness, however, we also reported the overall predicted count of cases for this age group from each model.

Extrapolation to the UK
Guided by our validation procedures, we identified which model had the greatest overall predictive validity, and proposed this as a candidate for FEP incidence prediction in England and Wales. We repeated out-of-sample prediction on the sociodemographic and socioenvironmental population characteristics of each LAD in England and Wales to obtain national-level and LAD-level predictions. Denominator data were obtained from the ONS 2009 mid-term estimates and stratified as previously described. Overall counts were derived for three broad age groups (16–35, 36–64 and 16–64 years), and for each of these, by sex and ethnicity. The 95% PIs were estimated as before. We visualised these data on maps and in tables to provide healthcare planners and commissioners with an easy-to-use tool to forecast the expected incidence of psychotic disorder in England and Wales. We have made this available as a free, open-use prediction tool, known as PsyMaptic (V0.5) (Psychiatric Mapping Translating Innovations into Care; http://www.psymaptic.org). Counts of cases predicted by our model were compared with those obtained under the Department of Health’s uniform rate in each LAD. We expressed these comparisons as ratios with 95% CIs derived using the same method as for standardised morbidity ratios (SMR). This approach was conservative because here we substituted the usual numerator in an SMR, the observed, O, for a predicted count. Unlike an observed count, no sampling variation is present for the predicted count, only uncertainty due to the model from which the prediction was estimated. Since variance in the prediction is therefore much smaller than the variance normally present for the numerator (O), this led to conservative estimates of 95% CI. Ratios in LAD where 95% CI did not span unity could therefore be interpreted as regions where there was strong evidence that the predictions from our model differed significantly from those predicted by the department of health’s uniform rate.

Software
All negative binomial regression models, out-of-sample prediction and estimation of 95% PI were conducted in R (V2.15.1). Cross-validation and model-fit diagnostics were conducted in Stata (V11). Prediction maps for England and Wales were created using StatPlanet Plus (V3.0) visualisation software.

RESULTS
Prediction sample
Our prediction models contained data on 1037 persons with a first episode psychosis in the ÆSOP (n=553; 53.3%) and ELFEP (n=484; 46.7%) studies, ascertained from over 2.4 m person-years at-risk. Twelve participants were excluded from the original ÆSOP sample because they were of no fixed abode and could not be geocoded to an LAD.

The population at-risk in the prediction sample came from LADs with higher median levels of multiple and employment deprivation, extent of deprivation and population density than the population at-risk in the validation sample, though there were no statistically significant differences in median income deprivation between the two samples (see online supplementary table S1).

Parameter coefficients obtained from the full prediction sample following negative binomial regression are shown in table 2. As previously reported from these data, incidence rates were generally raised in ethnic minority groups compared with the white British population. Models 2–6 included a measure of LAD deprivation (models 2–5) or population density (model 6), which were all significantly associated with an increased incidence of psychotic disorder, after control for individual-level confounders. Each of these models produced a lower AIC score than a model fitted solely with individual-level covariates (model 1), indicating a better fit. Cross-validation suggested that all models achieved good CCC agreement between predicted and observed cases, with low RMSE values (table 2).

Validation sample
Observed participants
We identified 572 potential participants over the first 30 months of the SEPEA study, aged 16–35 years, who...
### Table 2  Prediction models, covariates and fit: all clinically relevant psychoses (F10–39)

<table>
<thead>
<tr>
<th>Variable</th>
<th>Model 1 (IRR (95% CI))</th>
<th>Model 2 (IRR (95% CI))</th>
<th>Model 3 (IRR (95% CI))</th>
<th>Model 4 (IRR (95% CI))</th>
<th>Model 5 (IRR (95% CI))</th>
<th>Model 6 (IRR (95% CI))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age group×sex interaction*</td>
<td>p=0.07</td>
<td>p=0.06</td>
<td>p=0.06</td>
<td>p=0.06</td>
<td>p=0.06</td>
<td>p=0.06</td>
</tr>
<tr>
<td>Ethnicity</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>White British</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Non-British white</td>
<td>2.0 (1.5 to 2.5)</td>
<td>1.7 (1.4 to 2.2)</td>
<td>1.7 (1.3 to 2.2)</td>
<td>1.7 (1.3 to 2.2)</td>
<td>1.8 (1.4 to 2.3)</td>
<td>1.7 (1.3 to 2.2)</td>
</tr>
<tr>
<td>Black Caribbean</td>
<td>6.0 (4.9 to 7.3)</td>
<td>5.3 (4.3 to 6.5)</td>
<td>5.2 (4.3 to 6.4)</td>
<td>5.2 (4.3 to 6.4)</td>
<td>5.4 (4.5 to 6.6)</td>
<td>5.1 (4.2 to 6.3)</td>
</tr>
<tr>
<td>Black African</td>
<td>4.1 (3.3 to 5.1)</td>
<td>3.6 (2.9 to 4.5)</td>
<td>3.5 (2.8 to 4.4)</td>
<td>3.5 (2.8 to 4.4)</td>
<td>3.7 (3.0 to 4.6)</td>
<td>3.5 (2.8 to 4.3)</td>
</tr>
<tr>
<td>Indian</td>
<td>1.7 (1.2 to 2.5)</td>
<td>1.5 (1.1 to 2.2)</td>
<td>1.5 (1.0 to 2.2)</td>
<td>1.5 (1.0 to 2.1)</td>
<td>1.6 (1.1 to 2.3)</td>
<td>1.6 (1.1 to 2.2)</td>
</tr>
<tr>
<td>Pakistani</td>
<td>1.0 (1.2 to 2.5)</td>
<td>1.6 (1.0 to 2.4)</td>
<td>1.6 (1.0 to 2.4)</td>
<td>1.6 (1.0 to 2.4)</td>
<td>1.6 (1.1 to 2.5)</td>
<td>1.8 (1.3 to 2.7)</td>
</tr>
<tr>
<td>Bangladeshi</td>
<td>2.1 (1.5 to 2.8)</td>
<td>1.7 (1.2 to 2.3)</td>
<td>1.7 (1.2 to 2.3)</td>
<td>1.7 (1.2 to 2.2)</td>
<td>1.8 (1.3 to 2.5)</td>
<td>1.8 (1.4 to 2.4)</td>
</tr>
<tr>
<td>Mixed white and black</td>
<td>4.3 (2.8 to 6.7)</td>
<td>3.9 (2.5 to 6.0)</td>
<td>3.9 (2.5 to 6.0)</td>
<td>3.9 (2.5 to 6.0)</td>
<td>4.0 (2.8 to 6.1)</td>
<td>3.9 (2.5 to 6.1)</td>
</tr>
<tr>
<td>Caribbean</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixed, other ethnicities</td>
<td>1.3 (0.8 to 2.3)</td>
<td>1.2 (0.7 to 2.1)</td>
<td>1.2 (0.7 to 2.1)</td>
<td>1.2 (0.7 to 2.1)</td>
<td>1.2 (0.7 to 2.1)</td>
<td>1.2 (0.7 to 2.1)</td>
</tr>
<tr>
<td>Other ethnicities</td>
<td>2.2 (1.6 to 3.0)</td>
<td>1.9 (1.4 to 2.7)</td>
<td>1.9 (1.4 to 2.6)</td>
<td>1.9 (1.4 to 2.6)</td>
<td>2.0 (1.4 to 2.7)</td>
<td>1.9 (1.4 to 2.7)</td>
</tr>
</tbody>
</table>

Socioenvironmental variables

| IMD (z-score) | − | 1.184 (1.101 to 1.274) | − | − | − | − |
| Extent of deprivation (%) | − | − | 1.008 (1.004 to 1.011) | − | − | − |
| Income deprivation (%) | − | − | − | 1.025 (1.015 to 1.035) | − | − |
| Employment deprivation (%) | − | − | − | − | 1.062 (1.032 to 1.093) | − |
| Population density (pph) | − | − | − | − | 1.006 (1.003 to 1.007) | − |

Model-fit diagnostics

| AIC† | 2571.8 | 2552.4 | 2551.3 | 2549.6 | 2556.4 | 2556.3 |
| Mean Lin’s CCC (95% CI)‡ | 0.75 (0.74 to 0.77) | 0.77 (0.75 to 0.78) | 0.77 (0.75 to 0.79) | 0.77 (0.75 to 0.78) | 0.77 (0.75 to 0.78) | 0.76 (0.74 to 0.77) |
| Mean RMSE (SD)§ | 0.75 (0.11) | 0.74 (0.11) | 0.74 (0.10) | 0.74 (0.10) | 0.74 (0.11) | 0.76 (0.13) |

*All models fitted with age group by sex interaction given a priori evidence for effect modification. Likelihood ratio test p values reported between models with and without an interaction term fitted between age group and sex. Specific IRR has not been reported for clarity, but is available on request.

†Lower scores denote improved model fit.
‡Higher scores indicate greater correlation between observed and predicted count of cases in the prediction sample. Mean CCC and 95% CI reported following h=20 trials during cross-validation.
§Lower scores indicate lower prediction error. Mean RMSE and SD reported following h=20 repeats of K-fold cross-validation, where K=10.
AIC, Akaike Information Criterion; CCC, Lin’s correlation concordance coefficient; IRR, incidence rate ratio; RMSE, root mean squared error.
met initial acceptance criteria for EIS in East Anglia. We excluded 50 participants (8.7%) who did not meet clinical criteria for the ICD-10 psychotic disorder. This left an incidence sample of 522 participants from nearly 1.4 m person-years at-risk (37.4/100 000 person-years; 95% CI 34.3 to 40.7). A further 2.3 m person-years at-risk accrued in the same region for people aged 36–64 years over this period. Median levels of multiple, income and employment deprivation in the region did not differ significantly from the remainder of England, although the median population density and extent of deprivation in East Anglia were lower than elsewhere in England (see online supplementary table S1).

External model prediction and validation
The overall observed count of cases, aged 16–35 years, in the validation sample (n=522) fell within 95% PIs in four of seven models (models 3–6, table 3). Of these, the observed count (n=522) was closest to the point estimate for model 6 (508.5; 95% PI 449.0, 559.0), fitted with age group, sex, their interaction, ethnic group and LAD population density. The observed count of cases also fell within PIs from this model in five of six EIS in the study region, and 19 of 21 LADs, the most in any model (table 4). This model had the lowest error scores at the EIS (RMSE=11.6) and LAD (RMSE=6.1) levels of any model. Overall, model 6 was ranked highest across all external model-fit diagnostics (table 4). All models outperformed the department of health’s uniform figure of 51 per 100 000 person-years (model 7), which generally overestimated cases in the validation sample (overall prediction: 715.7 cases; 95% PI 664.0, 769.0).

We reported predicted cases aged 36–64 years from our models (table 4), although we could not test these in the validation sample. Model 6 predicted an additional 262.9 cases aged 36–64 years over a 2.5-year period in East Anglia (95% PI 233.0, 297.0).

We inspected the stratum-specific external validity of our best-fitting model (model 6, see online supplementary table S2), which performed accurately for sex-specific predictions, but less well in age-specific and ethnicity-specific strata. Thus, our model tended to underpredict observed cases in people aged 16–19 years, but overpredicted cases observed in people over 25 years old. With respect to ethnicity, model predictions were consistent with observed FEP cases for people of non-British white, black African, Bangladeshi and mixed ethnicities. However, our model tended to underpredict observed rates in the white British group, and overpredicted rates in the black Caribbean, Indian and Pakistani populations.

Extrapolation to England and Wales
We predicted the expected count and incidence of first episode psychosis per annum in each LAD in England and Wales based on model 6, and visualised these data in maps and tables freely available at www.psymaptic.org. Many maps can be visualised (eg, see online supplementary figure S1), including the overall predicted incidence counts and rates for each broad age group at the LAD level, and by sex. We will make PsyMaptic data available by ethnic group when we can improve the validity in ethnic-specific strata. According to our model, the annual number of new FEP cases in England and Wales would be 8745 (95% PI 8558, 8933), of which our model predicted 67.9% (n=5939; 95% PI 5785, 6102) would be seen through EIS. Only 176 (95% PI 151, 203) cases aged 16–64 years were forecast in Wales per annum. Assuming that our prediction model is accurate, it indicated that the Department of Health’s current uniform rate of 51/100 000 person-years was higher than the predicted point estimates for rates forecast by our PsyMaptic model in 351 LADs (93%) in England and Wales, but was lower than that predicted by our model in Birmingham and several London boroughs (see online supplementary figure S2, left-hand map). Under a conservative approach, these differences achieved statistical significance in parts of London (where the Department of Health’s model underestimated the need as predicted by PsyMaptic), and in some more rural parts of England and Wales (where the Department of Health’s model overestimated the need; see online supplementary figure S2, right-hand map).

DISCUSSION
Principal findings
We have developed and tested several epidemiological prediction models to forecast FEP incidence in England and Wales, having taken into account regional differences in the sociodemographic and socioenvironmental profiles of different populations. Inspection of our data suggested that a model fitted with age group, sex, their interaction, ethnic group and LAD-level population density provided the greatest external predictive validity when compared with the observed FEP caseload ascertained through EIS in our validation sample. This model also had good apparent validity across the entire age range (16–64 years). All models outperformed the Department of Health’s current gold standard for EIS commissioning, based on a uniform incidence rate. Our data suggested that the original figure used to commission EIS probably overestimated the true incidence of FEP in rural areas, and underestimated rates in urban settings. However, we acknowledge that commissioning decisions will need to be based on several additional factors, including the level of preclinical or non-psychotic psychopathology requiring assessment at initial referral to EIS, and variation in service organisation, remit and delivery.

Limitations and future development
Our prediction models were based on epidemiological data obtained from large, robust population-based FEP studies for people aged 16–64 years. The best-fitting model had good apparent validity over this age range,
## Table 3

<table>
<thead>
<tr>
<th>EIS</th>
<th>Observed</th>
<th>Predicted (95% PI) Model 1</th>
<th>Predicted (95% PI) Model 2</th>
<th>Predicted (95% PI) Model 3</th>
<th>Predicted (95% PI) Model 4</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Overall total, 16–35 years</td>
<td>522 (586.0, 696.1)</td>
<td>468.5 (422.0, 518.0)</td>
<td>474.7 (429.0, 522.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CAMEO North</td>
<td>84.7 (66.0, 106.0)</td>
<td>68.5 (52.0, 87.0)</td>
<td>67.8 (51.0, 86.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CAMEO South</td>
<td>163.6 (135.0, 192.0)</td>
<td>105.5 (84.0, 129.0)</td>
<td>111.7 (89.0, 134.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>West Norfolk</td>
<td>29.2 (18.0, 41.0)</td>
<td>23.0 (13.0, 35.0)</td>
<td>22.0 (12.0, 32.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Central Norfolk</td>
<td>162.6 (136.0, 191.0)</td>
<td>122.6 (99.0, 148.0)</td>
<td>123.0 (100.0, 149.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Great Yarmouth and Waveney</td>
<td>49.1 (34.0, 65.0)</td>
<td>41.6 (28.0, 55.0)</td>
<td>39.9 (27.0, 53.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Suffolk</td>
<td>151.9 (126.0, 178.0)</td>
<td>107.4 (85.0, 128.0)</td>
<td>110.3 (88.0, 133.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Overall total, 36–64 years</td>
<td>322.2 (292.0, 373.0)</td>
<td>244.0 (213.0, 276.0)</td>
<td>248.6 (216.0, 280.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Overall total, 16–35 years</td>
<td>477.1 (428.0, 523.0)</td>
<td>508.5 (459.0, 559.0)</td>
<td>715.6 (664.0, 769.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CAMEO North</td>
<td>69.1 (52.0, 88.0)</td>
<td>64.5 (48.0, 82.0)</td>
<td>92.1 (74.0, 111.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CAMEO South</td>
<td>100.7 (79.0, 123.0)</td>
<td>131.1 (108.0, 157.0)</td>
<td>169.0 (144.0, 195.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>West Norfolk</td>
<td>24.7 (16.0, 35.0)</td>
<td>22.3 (13.0, 33.0)</td>
<td>35.8 (25.0, 48.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Central Norfolk</td>
<td>128.2 (105.0, 153.0)</td>
<td>132.5 (108.0, 157.0)</td>
<td>187.7 (161.0, 215.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Great Yarmouth and Waveney</td>
<td>47.7 (35.0, 63.0)</td>
<td>38.0 (26.0, 51.0)</td>
<td>59.1 (44.0, 74.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Suffolk</td>
<td>106.6 (86.0, 128.0)</td>
<td>120.0 (96.0, 143.0)</td>
<td>172.1 (147.0, 198.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Overall total, 36–64 years</td>
<td>249.7 (221.0, 284.0)</td>
<td>262.9 (233.0, 297.0)</td>
<td>1175.4 (1109.0, 1243.0)</td>
</tr>
</tbody>
</table>

*Numbers in italics denote where the observed count fell within 95% prediction interval (95% PI) for people aged 16–35 years. Observed data for people aged 36–64 years in the validation sample were not available.

Model 1: Age group, sex, their interaction and ethnicity.
Model 2: Model 1+IMD.
Model 3: Model 1+extent of deprivation.
Model 4: Model 1+income deprivation.
Model 5: Model 1+employment deprivation.
Model 6: Model 1+population density.
Model 7: Department of health uniform figure for EIS of 15 new cases per 100,000 people/year.
and good external validity over the age range 16–35 years. While 16–35 years covers the majority of adult onset psychosis cases seen in mental health services, we recognise that some EIS teams incept people from 14 years old. We were unable to extrapolate our models to this age range, given the current absence of incidence data for this group in England. Data from Scandinavia suggest that the incidence of such ‘early onset’ psychoses is absolutely low, although the rate may have increased over the last few decades, probably as a result of movement towards earlier detection. We were also unable to externally validate prediction models for people aged 36–64 years, because comparable observed incidence data were not available in our validation sample. We have no reason to believe our predictions will be invalid for this group, however, since the empirical data which underpinned our models were ascertained from the same two large, well-conducted studies as for data on the younger age group. Furthermore, published findings from these studies are consistent with the wider epidemiological literature on FEP in England and internationally. It will be important to validate the predictive capability of our model(s) in this age range, and we will seek to identify suitable samples to do so in future versions of PsyMaptic.

Our best-fitting overall model demonstrated excellent external validity for predicting sex-specific FEP cases in our validation region (ie, SEPEA). It performed less well across age-specific and ethnic-specific stratum in this region. With respect to age, this discrepancy is most likely to be a function of EIS provision itself, which seeks to intervene as early as possible in the onset of psychosis. The effect of this will reduce median age at onset in comparison to studies conducted prior to the introduction of EIS, such as the ÅSOP and ELFEP studies upon which our models are based. Future versions of PsyMaptic will incorporate empirical data from post-EIS studies to improve age-specific predictions. The validity of our model in some ethnic groups also requires further refinement. Much of the prediction data underlying our models came from urban environments with large proportions of ethnic minority groups. The sociodemographic profile and sociocultural experiences of these groups may be very different to those of their counterparts in other, less urban, parts of England, thus altering psychosis risk in different ethnic groups. In our observed data, a larger proportion of cases were white British than predicted by our model. If ethnicity is a partial proxy for exposure to deleterious socioeconomic experiences, such as the combined effect of social inequality, fragmentation, deprivation and population density, then simultaneously incorporating such factors into our models may improve their predictive validity by ethnicity. Alternatively, risk by ethnic group may be conditional upon (ie, interact with) environmental factors in urban areas (as with the ethnic density effect), but whether such interactions exist in less urban regions is not known. Forthcoming SEPEA and PsyMaptic data will explore such possibilities.

All prediction models had reasonable apparent validity, although our proposed model performed slightly worse (most noticeably for AIC) than models which included deprivation (ie, models 2–4) instead of population density. Our decision to use model 6 as our proposed candidate for the prediction tool was supported by the fact that it produced the most accurate external forecasts of any model, despite considerable socioenvironmental differences between regions in our prediction and validation samples. We were unable to predict the expected incidence of psychotic disorder in geographical areas smaller than LADs, such as electoral wards, or to other parts of the UK, because appropriate denominator data were not published as mid-term census estimates. The 2011 census will provide small area and national data for the whole of the UK, scheduled for release in mid-2013. This will allow us to update our tool to the latest population estimates for the UK, and refine our PsyMaptic tool at a smaller geographical level for fine-grained healthcare commissioning. We will then be able to develop models to explore cross-level interactions, such as the association between individual ethnicity and neighbourhood-level ethnic density. Small

<table>
<thead>
<tr>
<th>Table 4 External model validation diagnostics*</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Observed case count within SEPEA overall prediction intervals? (rank)</strong></td>
</tr>
<tr>
<td>Number correct (rank)</td>
</tr>
<tr>
<td>Model 1 No (5)</td>
</tr>
<tr>
<td>Model 2 No (5)</td>
</tr>
<tr>
<td>Model 3 Yes (1)</td>
</tr>
<tr>
<td>Model 4 Yes (1)</td>
</tr>
<tr>
<td>Model 5 Yes (1)</td>
</tr>
<tr>
<td>Model 6 Yes (1)</td>
</tr>
<tr>
<td>Model 7 No (5)</td>
</tr>
</tbody>
</table>

*For each diagnostic, models are placed in rank order (1=best model, 7=worst model) with ties given the same ranking. The mean ranking and rank provide an estimate of the overall performance of various models.

EIS, early intervention services; LAD, local authority district; RMSE, root mean squared error; SEPEA, Social Epidemiology of Psychoses in East Anglia.
area prediction models will require a multilevel approach, not attempted here, because obtaining predictions from multilevel random effects models is not straightforward and requires active statistical development.

We believe case ascertainment in our validation sample led to a reliable estimate of the incidence of psychotic disorder for people aged 16–35 years. EIS were the only mental health service for people aged 14–35 years experiencing a first episode of psychosis in East Anglia, minimising the potential for underascertainment in the population at-risk when derived from careful epidemiological design.\(^{13}\) We are confident that our validation sample also contained few false positive cases for any clinically relevant psychoses, since participants were excluded who failed to meet acceptance criteria for EIS, or who did not meet clinical diagnosis for psychotic disorder in the first 6 months following EIS acceptance. It is important to recognise that while our prediction models are based on diagnosed clinically relevant psychoses, service commissioning will also need to account for additional preclinical or non-psychotic psychiatric morbidity presenting to EIS, particularly in services which operate early detection models or implement ‘watch-and-wait’ briefs. The SEPEA data used to validate our models do not predict (1) the number of ‘false positive’ subjects who may require psychiatric triage and assessment, even though they are not accepted by EIS or (2) the number of ‘true positive’ subjects accepted by services, but who did not meet epidemiological criteria for inclusion in the validation sample of the SEPEA study (ie those living outside the catchment area at first contact, or those transferred from other services); these people will consume varying degrees of service resources which need to be considered in service planning.

We also note that pathways to care may affect the level of incidence observed in EIS, since many filters are likely to operate before subjects come to the attention of EIS. These will include local level service organisation and the relationship between Community Mental Health Teams, Child and Adolescent Mental Health and EIS. Furthermore, acceptance criteria for entry to EIS vary, which will have a downstream effect on the number of new cases of clinically relevant psychoses received in each team. Future versions of PsyMaptic will include forecasts for specific psychotic disorders, as standardised research-based diagnoses (using OPCRIT\(^{34}\)) are currently being collected in the ongoing SEPEA study. Acceptance rates to EIS may also be influenced by local community awareness of such services. While our prediction models outperformed the current gold standard for EIS commissioning in England when restricted to clinically relevant caseloads, we recommend that our models are best interpreted as forecasts of the expected burden of first episode psychosis in given populations, not the total burden of resource consumption through EIS, given these issues.

We estimated PIs from first principles (DJ) since their derivation is an area of statistical development.\(^{35}\) We used a bootstrap-like methodology to produce 95% PI accounting for natural variation in the validation sample, but ignoring parameter uncertainty in the coefficients included in prediction models, which we assumed to be the true coefficients of risk in the population. Our approach therefore naturally led to slightly artificially narrow 95% PIs. This was not necessarily undesirable for the purpose of model validation and the precise prediction of expected counts, because we wished to apply stringent criteria. Ideally, 95% PIs should take into account both these sources of variation, although we note that parameter uncertainty is usually small compared with the natural variation of the quantities of interest. The addition of more empirical data in the prediction sample would not lead to narrower 95% PIs, though it would tend to move the point estimate of risk for each coefficient closer to the true value in the population. We do not believe we have misestimated the point estimates of risk across major sociodemographic groups, since our results accord with the wider literature.\(^{17,21,22}\)

We sought independent confirmation that our development of 95% PI was correct (personal communication with Professor Ian White, MRC Biostatistics Unit). We recommend that all prediction point estimates from our PsyMaptic model are considered with their 95% PIs, which provide information about the natural variance in expected rates in the population.

**Meaning of the findings**

If commissioners are to meet the Department of Health’s vision to orientate health services around local need,\(^1\ 2\ 5\) differences in the demand for EIS and other mental and physical health services will need to be taken into account to allocate finite resources where they are most needed. The PsyMaptic prediction model provides proof-of-concept that when robust empirical epidemiological data are combined with accurate population at-risk estimates, this can be realised. As such, our modelling approach could have utility in many other settings and for many disorders. Our translational approach demonstrated good validity to predict the expected incidence of first episode psychosis, particularly through EIS, where 76% and 63% of all male and female adult-onset FEP cases, respectively, will typically present.\(^{38}\) Since their inception in 2002, EIS in England and Wales have reported both lower\(^{11}\) and higher\(^{16}\) caseloads than they were originally envisioned to manage,\(^5\) with shortfalls or excesses in anticipated demand for services aligned to the degree of urbanisation in the underlying catchment area. Others have noted that EIS provision in rural areas may be difficult to implement effectively,\(^{14}\ 15\) and while the MH-PIG acknowledged that “...an understanding of local epidemiology is needed as the size of population covered will depend on a number of different factors” (ref. 5, p. 55), no further elaboration on how to achieve this was provided. We
believe PsyMaptic provides a possible tool to overcome this challenge, improving the description and prediction of local population need beyond the MH-PIG and including individual-level and neighbourhood-level indicators of local need.\textsuperscript{17} From an aetiological perspective, we acknowledge that variables such as ethnicity or population density are likely to be markers for a suite of more complex, interactive social, genetic and environmental determinants of psychosis.\textsuperscript{36}

Our models are not the first to be used to forecast mental illness needs in England and Wales,\textsuperscript{37} though we believe this is the first attempt to forecast incidence rather than prevalence in the community. We recommend that our prediction methodology is used in conjunction with demand for services as predicted by PsyMaptic.\textsuperscript{41}

Ongoing monitoring and audit of EIS will be vital to ensure that services meet the fidelity criteria upon which they were originally commissioned,\textsuperscript{11} \textsuperscript{40} including ensuring that service capacity matches local need as closely as possible. As part of this process, we will need to externally validate our models in a wider range of settings, refining them based on empirical observation.

We note that advocacy expressed for EIS by healthcare professionals in England and Wales broadly correlates with demand for services as predicted by PsyMaptic.\textsuperscript{41} Though by no means universal, proponents of EIS tend to be located in major conurbations—such as London,\textsuperscript{42} Birmingham\textsuperscript{43} or Manchester\textsuperscript{44}—where the demand for EIS will be highest, while those who suggest EIS resources could be used more effectively through other types of mental health service provision tend to work in more rural communities,\textsuperscript{15} \textsuperscript{41} where but a handful of young people would be expected to come to the attention of services each year. It is possible that both sides are correct and that more resources are required to help with the tide of psychotic illness in inner cities. Resources might be used more effectively in other ways, elsewhere, so long as the needs of the small number of young people who suffer an FEP each year are met; a dedicated specialist EIS may not be the most effective approach when anticipated demand will be very low.

Given the significant downstream economic savings associated with spending on EIS as estimated in an urban setting,\textsuperscript{8} PsyMaptic could be used to highlight regions where sufficient investment to appropriate mental health services would lead to the greatest economic gains in terms of mental healthcare expenditure (assuming sustained intervention also leads to improved social and clinical benefit for patients\textsuperscript{6} \textsuperscript{7}). PsyMaptic can also be used to highlight regional variation in demand according to age and sex and, in future versions, by ethnicity. This will allow service planners to tailor provision around the sociocultural characteristics of their local populations. Our prediction tool for first episode psychosis, which translates robust empirical epidemiological data on psychosis risk to the population structure of different regions, offers a methodology for improving the allocation of finite mental health resources based on local need.

Author affiliations
\textsuperscript{1}Department of Psychiatry, University of Cambridge, Herchel Smith Building for Brain & Mind Sciences, Cambridge, UK
\textsuperscript{2}MRC Biostatistics Unit, Institute of Public Health, University of Cambridge, Forvie Site, Robinson Way, Cambridge, UK
\textsuperscript{3}CAMEO, Cambridgeshire & Peterborough NHS Foundation Trust, Cambridge, UK
\textsuperscript{4}Norfolk and Suffolk Partnership Trust, Ipswich Hospital, Norwich, UK
\textsuperscript{5}Suffolk Early Intervention Psychosis Service, Norfolk and Suffolk Partnership Trust, Stowmarket, Suffolk, UK
\textsuperscript{6}Forensic Psychiatry Research Unit, Queen Mary’s University London, St. Bartholomew’s Hospital, London, UK
\textsuperscript{7}Department of Psychiatry Studies, Institute of Psychiatry, London, UK
\textsuperscript{8}NIHR Collaboration for Leadership in Applied Health Research & Care, Cambridge, UK

Acknowledgements The authors are grateful to the clinical services and staff participating in the SEPEA study, and the MHRN for their support. We are grateful to Professor Ian White of the MRC Biostatistics Unit (University of Cambridge) for his guidance on developing our prediction interval methodology for negative binomial regression.

Contributors JKB was responsible for the concept, design, analysis, data extrapolation, interpretation of the data, as well as for drafting the report and developing the content for the websites www.psymaptic.org, www.psymaptic.com and www.psymaptic.co.uk. He was also the chief investigator of the SEPEA study, where the validation sample data were obtained. JKB gave final approval of this version of the manuscript to be published. DJ was responsible for developing and initially implementing the statistical approach with regard to prediction intervals from count-based regression. He also contributed to drafting and editing the final version of this manuscript. He gave final approval of this version of the manuscript to be published. JPB is the principal collaborator on the SEPEA study in the CAMEO early intervention in psychosis services. He also contributed to editing the final version of this manuscript. He gave final approval of this version of the manuscript to be published. DJ was the principal collaborator on the SEPEA study in the CAMEO early intervention in psychosis services. He also contributed to editing the final version of this manuscript. He gave final approval of this version of the manuscript to be published. FPW is the principal collaborator on the SEPEA study in the CAMEO early intervention in psychosis services. He also contributed to editing the final version of this manuscript. He gave final approval of this version of the manuscript to be published. JC is the principal investigator of the ELFEP study and provided part of the empirical dataset underlying the prediction model for use in this project. He also contributed to editing the final version of this manuscript. He gave final approval of this version of the manuscript to be published. PSJ is the principal collaborator of the SEPEA study in the CAMEO early intervention in psychosis services. He also contributed to editing the final version of this manuscript. He gave final approval of this version of the manuscript to be published. FPW is the principal collaborator of the SEPEA study in the CAMEO early intervention in psychosis services. He also contributed to editing the final version of this manuscript. He gave final approval of this version of the manuscript to be published. JPB is the principal investigator of the SEPEA study in the CAMEO early intervention in psychosis services. He also contributed to editing the final version of this manuscript. He gave final approval of this version of the manuscript to be published. FPW is the principal collaborator of the SEPEA study in the CAMEO early intervention in psychosis services. He also contributed to editing the final version of this manuscript. He gave final approval of this version of the manuscript to be published.

Funding Wellcome Trust (grant number WT085540) and NIHR (grant RP-PG-0606-1335).

Competing interests All authors declare that JKB has support from the Wellcome Trust for the submitted work (grant number WT085540) and PBJ has support from the NIHR (grant RP-PG-0606-1335).

Psychosis incidence prediction

Ethics approval
Ethics approval to conduct the original AESOP and East London First Episode Psychosis studies was granted from local research ethics committees in their respective centres prior to the original date of onset of the study. We were granted ethical approval to conduct the work related to the present manuscript, including the use of data from the SEPEA study, from the Cambridgeshire 3 REC (09/H0306/39).

Provenance and peer review
Not commissioned; externally peer reviewed.

Data sharing statement
Extra data are available at our prediction website, PsyMaptic: http://www.psymaptic.org. We have made a large array of prediction data freely available to interested users, who can visualise prediction data in maps and tables for a range of psychotic disorder predictions by major sociodemographic subgroups and by LAD. Future versions of PsyMaptic will release data for other regions and at other geographical levels. Users can also download LAD-specific and national summaries of prediction data from our models in tabular form. Finally, additional explanatory material about how to use the data and the PsyMaptic website is also available at http://www.psymaptic.org. Bespoke data not provided freely on our website may be requested from the authors at the cost of data extraction. The empirical data underlying the prediction models (AESOP and ELFEP) or the validation sample (SEPEA) are not freely available as the studies are ongoing, but papers detailing the main findings from these studies have previously been published elsewhere.

REFERENCES

A population-level prediction tool for the incidence of first-episode psychosis: translational epidemiology based on cross-sectional data

James B Kirkbride, Daniel Jackson, Jesus Perez, David Fowler, Francis Winton, Jeremy W Coid, Robin M Murray and Peter B Jones

BMJ Open 2013 3:
doi: 10.1136/bmjopen-2012-001998

Updated information and services can be found at:
http://bmjopen.bmj.com/content/3/2/e001998

These include:

Supplementary Material
Supplementary material can be found at:
http://bmjopen.bmj.com/content/suppl/2013/02/02/bmjopen-2012-001998.DC1.html

References
This article cites 32 articles, 9 of which you can access for free at:
http://bmjopen.bmj.com/content/3/2/e001998#BIBL

Open Access
This is an open-access article distributed under the terms of the Creative Commons Attribution Non-commercial License, which permits use, distribution, and reproduction in any medium, provided the original work is properly cited, the use is non commercial and is otherwise in compliance with the license. See: http://creativecommons.org/licenses/by-nc/2.0/ and http://creativecommons.org/licenses/by-nc/2.0/legalcode.

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Topic Collections
Articles on similar topics can be found in the following collections

Epidemiology (1626)
Evidence based practice (538)
Health policy (513)
Health services research (1067)
Mental health (512)
Public health (1680)

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/